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2,832,750

ORGANO TIN COMPOUNDS, METHOD OF MAK-ING SAME, AND MIXTURES THEREOF WITH HALOGEN-CONTAINING RESINS

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12 Claims, (CL 260-45.75)

The present application is a continuation in part of copending application Serial No. 232,878, filed June 21, 1951, now Patent No. 2,648,650, and relates to an invention pertaining to novel organotin derivatives of mercapto compounds and to resin compositions containing these compounds as a stabilizer.

In the aforesaid copending application, there is described and claimed broadly a new group of compounds generally designated as condensation products of certain organotin compounds with mercapto acids and esters. More particularly these compounds may be illustrated by 25 the following structural formulas:

R₂SnSR'COOR" R₂Sn(SR'COOR")₂ and RSn(SR'COOR")₃

wherein R is a univalent organic radical such as an alkyl, aryl, aralkyl or alkaryl, R" represents hydrogen or a univalent organic radical such as alkyl, aryl, aralkyl or alkaryl, and R' is a divalent organic radical selected from the class consisting of substituted or unsubstituted alkylene, arylene, aralkylene or alkarylene. It is to be noted that the tin radical is bonded to the sulfur radical.

The organo-tin derivatives referred to are far superior stabilizers for resins, particularly vinyl halide resins, to any known stabilizer in quantities as low as one half of one percent of the weight of the resin. This superior stabilizing property is equally applicable to resin-plasticizer compositions, inclusive of the phosphate type plasticizers, the use of which has heretofore been phohibitive due to their instability even in the presence of known stabilizers. However, the mercapto acid ester derivatives and the like of organo-tin compounds impart stabilization to the phosphate plasticizer resin systems, permitting thereby the production of superior fireproof plastic compositions.

These compounds may be prepared in any suitable manner. It has been found however that these novel products may be prepared in high yield and purity by reacting an organo-tin compound with mercapto-acids and mercapto-acid esters, and recovering a reaction product having the desired So-S linkage and ratio. More specifically, an organo-tin compound selected from the group consisting or organo-tin hydroxide, oxide, and stannoic acid may be condensed with an appropriate 60 amount of a mercapto acid or mercapto acid ester to produce products having the above structural formulas. An organo-tin halide having the formula RSnX, RsSnX, or R₂SnX wherein X is a halide, such as chlorine, may also be employed as the reactant with the mercapto acids 65 or esters. H wever, for optimum results, it is preferable to conduct these reactions in the presence of basic substances or alkaline neutralizing agents (hydrogen ion acceptors) such as oxides, hydroxides (e. g. sodium hydroxide) carbonates (e. g. sodium and potassium carbonate), and tertiary amines (e. g. pyridine), etc. These substances accept the hydrogen ion formed during the

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condensation reaction and are not deleterious to the mercapto acid or ester when used in the prescribed manner. Basic substances of the character of ammonia, primary and secondary amines, should not be employed ordinarily, because they tend to react with esters, and thereby decrease the yield of desired ester products.

The reaction mechanism for the formation of the bismercapto derivatives using organo-tin oxide is clearly illustrated by the following equation:

R₂SnO+2HSR'COOR"=R₂Sn(SR'COOR")_x+H₂O

This equation graphically shows that the reaction occurs between the mercapto radical and the organo-tin oxide, whereby water splits off and a chemical bond is formed between the tin and the sulfur. It further illustrates the necessity of two molecules of the mercapto compound per mole of tin containing reactant in order to obtain the desired reaction and the desired end product having a 2:1 ratio of sulfur to tin.

Similarly, products of the formula RSn(SR'COOR")₂ require a 3:1 ratio of sulfur to tin. Using stannoic acid as a reactant, the reaction may be illustrated as follows:

RSnOOH+3HSR'COOR"=RSn(SR'COOR")3+2H3O

Products of the formula R₂SnSR'COOR" require a 1:1 sulfur to tin ratio. Thus using organo tin chloride as a reactant, the reaction may be graphically represented as follows:

R_SnCI+HSR'COOR"=R_SnSR'COOR"+HCI

It is within the scope of this invention that pure, impure or commercial grades of the reactants may be employed satisfactorily. In general, pure compounds of the above formulas may be prepared from pure raw materials. However, these novel compounds may be diluted with innocuous, inert materials, thereby permitting the use of technical materials or intermediates in their preparation.

Any suitable reaction temperatures may be employed. It is ordinarily preferred to use room or slightly elevated temperatures of the order of up to about 50° C. The exothermic nature of the reaction between the organo-tin reactants and alpha or beta mercaptoacid esters and the like, affords a considerable saving in the expenditure for external heat. This reaction may be conducted at room temperature with a concomitant rise of about 5 to 10 degrees after the reaction has started. Thus, for example, the initial reaction temperature may be within the broad range of 19° C.-28° C., and the final reaction temperature may be within the ranges of 24° C.-40° C. However, the presence of the water produced by the condensation reaction usually requires an additional heating or refinxing in order to strip said water from the reaction product, though it may be removed in any suitable manner.

The use of inert organic solvents as the medium for the reaction, such as toluene, benzene, methyl alcohol, etc. is contemplated. The presence of such solvent facilitates the desired reaction. The solvent may be eliminated from the reaction product at the completion of the reaction by any suitable means. This may be accomplished by vaporizing the solvent under vacuum at elevated temperatures. Pressures of about 2 to 30 mm, and temperatures of 75°-160° C. are satisfactory in effecting removal of toluene or like solvent, from the reaction product.

By the procedures of this invention, these novel organotin derivatives of mercapto-acids and mercapto-acid o esters can be obtained in almost theoretical yields. These yields are indicative of the non-necessity of the use of an excess f either reactant, the total amount of starting 3

materials being substantially completely utilized in the formation of the final reaction product.

The following examples are further illustrative of these novel products and their preparation, and it will be understood that the invention is not limited thereto.

EXAMPLE 1

Mercaptoacetic acid, 97% (194 gms.), 3,5,5 trimethyl hexyl alcohol (288 gms.), paratoleene-sulphonic acid (2 gms.) (esterification catalyst) and 100 ml. of toluene were 10 placed in a glass reaction vessel provided with a Barrett moisture trap, glass agitator and immersion thermometer. The mixture was refluxed with stirring to remove water of reaction. In 6 hours, theoretical water was extracted. The final pot temperature was 160° C. The 15 toluene was then stripped in vacuo at 10 mm. pressure; the trimethyl hexyl mercaptoacetate was distilled at 125° C. at 10 mm. Sulfur content as calculated was 14.68% and as actually found was 14.68%.

The trimethyl hexyl mercaptoacetate (333 gms.) was added to a stirred slurry of 189 gms. dibutyltin oxide in 350 cc. of toluene. An exothermic reaction in which the temperature rose from 28° C. to 38.5° C. crsued. The slurry became slightly turbid in contrast with the initial opaque appearance. The temperature was then raised to toluene reflux. Using a moisture trap, water (11.8 gms.) was removed from the product. Toluene was removed by vacuum distillation: The dibutyltin S,S' bis(3,5,5 trimethylhexyl mercaptoacetate) was a slightly viscous colorless liquid (98% yield). The calculated analysis would be Sn, 17.76% and S, 9.62%. Analysis showed Sn, 17.41% and S, 9.36%.

EXAMPLE II

Cyclohexanol (150 gms.), mercaptoacetic acid (97%—146 gms.) and 50 ml. of toluene were placed in a reaction flask, fitted with a Barrett moisture trap and reflux condenser. The theoretical amount of water (32 cc.) was separated in 1½ hours. The final pot temperature was 160° C. The resulting cyclohexyl mercaptoacetate distilled at 105–10° C. at 10 mm. pressure. Sulfur content calculated 18.40%; found 18.11%.

The cyclohexyl mercaptoacetate (179 gms.) was added to 126 gms. of dibutyltin oxide slurried in 240 cc. of toluene. An exothermic reaction ensued—the temperature rising from 30° to 40° C. and, at the same time, the mixture became clearer. The mixture was then heated to remove water of reaction via a Barrett moisture trap (8.8 cc. were separated). The final pot temperature was 119° C. The toluene was then stripped at 18 mm. pressure; the final pot temperature was 135° C. The dibutyltin S,S' bis (cyclohexyl mercaptoacetate) was a slightly yellow liquid which had the following analyses: Sn=19.91%; S=10.80%. A calculated composition calls for Sn=20.30%; S=10.93%.

EXAMPLE III

Phenoxyethanol (276.4 gms. Dowanol 1), mercapto-acetic acid, 97% (200 gms.) and 100 ml. of toluene were refluxed and the theoretical amount of water was removed in 3 hours. The toluene and excess mercaptoncetic acid were stripped by heating to 100° C. at 10 mm. Sulfur content found 12.77%; calculated 15.10%.

The phenoxyethyl mercaptoacetnte (167 gms.) was added to 83 gms. of dibutyltin oxide slurried in 260 cc. of toluene. Temperature rise from 23° to 28° C. was noted. 5.1 cc. of water were removed in one hour. After the solution was clarified by filtration, the toluene was stripped to a pot temperature of 125° C. at 9 mm. pressure. The dibutyltin S,S' bis (phenoxyethyl mercaptosure. The dibutyltin S,S' bis (phenoxyethyl mercaptosure.) was a light colored liquid. It analyzed 14.10 Sn and 9.51% S. Calculated Sn and S contents are 16.23% and 8.73% respectively. Due to the technical nature of D wanol 1, the ester and reaction product are contaminated with inert diluents.

EXAMPLE IV

Dihydroahietyl mercaptoacetate was prepared by refluxing a toluene solution of ½ mole of "Abitol" (85% hydroahietyl alcohols, 166 gms.) with ½ mole of mercaptoacetic acid (48 gms.). About 9 cc. of water were removed at the time the esterification was stopped. No catalyst was used.

To this toluene solution of the hydroabietyl mercaptoacetate was added dibutyltin oxide (62.5 gms.—14 mole). The exothermic reaction which resulted caused the temperature to rise from 22° C. to 28° C. An hour of refluxing removed the water of reaction (4.5 cc.). The solution was filtered and the toluene was stripped in vacuo. The dibutyltin S,5° bis(dihydroabietyl mercaptoacetate) analyzed 11.05% Sn, 5.68% S.

EXAMPLE V

Trimethyl nonyl mercaptoacetate was prepared by placing 1 mole, 95 gms., of 97% mercaptoacetic acid and 1 mole, 186 gms., of commercial trimethyl nonyl alcohol with 100 ml. of toluene and boiling chips in a reaction flask. No esterification catalyst was used. The theoretical amount of water was over in about 14 hours. This solution of the ester was then reacted with 124.5 gms., 1/2 mole, of dibutyltinoxide. An exothermic reaction ensued in which the temperature rose from 22° C. to 31.5° C. 7.8 cc. of water was removed in two hours of refluxing. This toluene solution was filtered by gravity. The toluene was stripped from the batch at 25 mm. pressure to a pot temperature of 80° C. The pressure was then reduced to 8 mm. and the pot temperature was raised to 120° C. The residue, dibutyltin S.S. bis (trimethyl-nonyl mercaptoscotate), was a cloudy yellow viscous liquid, analysis of which was 12.5% Sn and 7.7% S. Various diluent impurities are present in the product due to the commercial nature of the reactants.

EXAMPLE VI

Diethylene glycol laurate mercaptoacetate was prepared from commercial diethylene glycol monolaurate. An estimated 1 mole of this latter material, 288 gms. was reacted with 95 gms. of mercaptoacetic acid (97%) in the presence of 50 ml. of toluene. The reaction was completed in 5 hours. This toluene solution was further reacted by adding 124.5 gms. of dibutyltin oxide to it. There was an exothermic reaction in which the temperature rose from 23° C. to 29° C. The reaction was completed in about 2½ hours wherein 7 cc. of water were removed. The toluene solution was filtered hot by gravity. The clear solution was then stripped to a final pot temperature of 100° C. at 10 mm. pressure. The resulting product, dibutyltin S,S' bis(diethylene glycol laurate mercaptoacetate) analyzed 9.4% Sn and 5.6% S.

EXAMPLE VII

Ethylene glycol ricinoleate mercaptoacetate was prepared in a manner similar to that used in Example VI. Since the ethylene glycol ricinoleate was a commercial product, the resulting materials were not of a very high purity. The ester so prepared was reacted by adding it to the toluene suspension of the dibutyltin oxide, ¼ mole, 62.5 gms. An exothermic reaction resulted in which the temperature rose from 23° C. to about 30° C. The theoretical water was removed in about one hour. The solution was filtered and the toluene was stripped. The resulting material was a thick tan liquid in which considerable precipitation resulted. The material analyzed 5.8% Sn and 3.8% S.

EXAMPLE VIII

2-butyloctanol-1, 176 gms., and beta mercaptopropionic acid, 100 gms., and 100 cc. of toluene were placed in a reaction flask and refluxed until 16 cc. of water were removed. This esterification proceeded at a somewhat alower rate than the mercapto-acetic acid esterifica-

tion. The toluene was stripped from this ester. The ester analyzed 10.6 S. The ester, 150 gms., was dissolved in 200 cc. of toloroe. 62.5 gms. of dibutyttin oxide were added to the tolnene solution and the temperature rose from 19° to 24° C. Toluene was refuned to remove water and 4.5 cc. were taken off in 3 hours. The toluene solution was filtered. The toluene was stripped from the solution at a pressure of 9 mm. to a pot temperature of 125°. The dibutyltin S.S' bis (2-butyloctyl beta mercaptopropionate) was a viscous slightly yellow liquid. 10 It analyzed 14.2% Sn. 7.5% S. Calculated analyzes, 14.0% Sn. 8.3% S.

EXAMPLE IX

captoacetic acid, 97% (95 gms.) with butyl alcohol (148 gms.) in the presence of paratolnene-sulfonic acid and removing water by means of a moisture trap. The excess alcohol was removed by vacuum distillation. ester was distilled at 80° C. at 20 mm. pressure. Sulfar 20 content: found 21.04; calculated 21.6%. 37 gms. of the butyl mercaptoacetate was mixed with 59.1 gms. of dilauryltin oxide using 100 ml. of toluene as the solvent. This slurry was refluxed until 214 ml of water were removed. The toluene solution was filtered. The toluene 25 was stripped to a pot temperature of 130° C at 2½ mm. A yellow liquid was obtained weighing 87.1 gms. The dilauryhin S,S' his (butyl mercaptoscetate) analyzed 15.2% Sn, 8.7% S. Calculated analyses 15.8% Sn, 8.5% S.

EXAMPLE X

Mixed decyl mercaptoacetate was prepared by reacting 1215 gms. of mercaptoacetic acid with 2000 gms, of mixed decanola (product of Carbide and Carbon Chemicals Corp.) in the presence of 12 gms. of para toluenesulphonic acid. In four hours 240 mls, of water were removed. The toluene was stripped in vacuo and the mixed decyl mercaptoacetates distilled at about 130° C. at 10 mm. pressure. The decyl mercaptoscetate, 62.5 gms., 81.3 gms. tributyltin monochloride, and 100 ml. of toluene were charged into a flask. To this flask was added gradually sodium carbonate, 26.5 gms., 100% in excess of the amount required. On adding the sodium carbonate, the temperature rose from 22° C. to 28° C. After addition, the mixture had a pink coloration. Heat was then applied. The reaction was completed in six hours as evidenced by the removal of water and carbon dioxide from the reaction. The salt residue was filtered and this material was stripped of toluene via vacuum 50 distillation to a pot temperature of 125° C. at 2 mm. pressure. The product was a yellow colored liquid which weighed 107 grams. The tributyltin S decyl mercaptoacetate analyzed 21.3% Sn, 6.7% S. Theoretical analyear for such a material is 22.8% Sn, 6.2% S.

EXAMPLE XI

Decyl mercaptoacetate, 35.8 gms. of a 93% purity, was admixed with butyl stannoic acid, 9.8 gms., and 50 ml. of toluene. The mixture was heated with stirring and water was removed as it was formed. The reaction was completed in 21/2 hours. The solution was filtered by gravity and the toluene was stripped in vacuo to a pot temperature of 140° C. at 2 mm. The product was a light yellow liquid weighing 36.7 gms. The product analyzed 13.25% Sn. 11.1% S. The theoretical analysis for this product which is butylin S.S'.S" tris(decyl mercaptoacetate) is 13.7% Sn. 11.0% S.

EXAMPLE XII

Decyl mercapioacetate, 14.7 gms. of a 93% purity, was 70 admixed with triphenyltin chloride, 22.7 gms., and 55 ml of toluene. To this mixture was added slowly sodium carbonate, 10.6 gms. The addition of the carbonate and removal of water of reaction were completed in 31/2 hours. The material was then filtered by gravity and the 75

tohiene was stripped in vacuo to a pot temperature of 130° C. at 2 mm. pressure. The product was a light colored viscous liquid. It analyzed 19.4 Sn. 6.1 S. The theoretical analysis for this triphenyltin S decyl mercaptoacetate is 20.5 Sn. 5.5

EXAMPLE XIII

Thiomalic acid (87.0 gms.—0.50 mole) was dissolved in methyl alcohol (300 ml.). The solution was stirred while dibutyltin oxide (62.5 gms.—0.25 mole) was added. The exide dissolved. The reaction was completed by distilling off the methyl alcohol and water of reaction in vacuo at a pot temperature of 105° C. and then placing the reaction product on a steam bath in an evaporating Butyl mercaptoaccuate was prepared by refluxing mer- 15 dish. The product, dibutylin S.S' his(thiomalic acid) ptoacctic acid, 97% (95 gms.) with butyl alcohol (148 was a grey, low-melting solid which analyzed 18.9% Sn. 11.3% S.

EXAMPLE XIV

Part A

Three moles (325.5 grams) of alpha chloropropionic acid were added dropwise to a solution of sodium milftydrate (6 moles, 480 grams of 70% sodium sulfhydrate and 250 ml. of water) saturated with hydrogen sulfide. The mixture was stirred and hydrogen sulfide was passed into the solution during the addition. After the acid had been added the solution was heated to reflux for 31/4

The solution was then cooled and an excess of hydrochloric acid was added. The solution was heated for three hours to effect complete reaction. It was then cooled and extracted with other.

The other extract was evaporated until free of other and the residue distilled to recover the mercapto-acid. Yield was 90 grams of a clear liquid.

Two moles of 3,5,5 trimethylhexanol, the 90 grams of alpha mercaptopropionic acid, 100 ml. of toluene and 2 ml. of 47% boron trifluoride etherate were refluxed until the water of esterification had been removed. To the resulting ester, 0.42 mole (104.5 grams) of dibutyltin oxide was added, the mixture refluxed to effect solution of the oxide and removal of water of reaction. The final mixture was stripped under vacuum to leave as a residue dibutyltin S,S' bis(trimethylhexyl e-mercaptopro-

EXAMPLE XV

The process of Example XIV was repeated using 1 mole of \$-bromobutyric acid in place of the e-chlorobutyric acid. The other ingredients were used in equivalent amounts. The resulting product is dibutyitin S.S. bis (trimethylhexyl β-mercaptobutyrate).

EXAMPLE XVI

Three moles of e-bromobutyric acid and 3.15 moles of thioures were heated in 500 ml. of alcohol to form 5-ethylpseudothiohydantoin hydrobromide. The pseudothiohydantoin was isolated by evaporation of the alcohol and hydrolyzed by boiling sodium hydroxide solution in alcohol. This method is reported in J. Am. Chem. Soc. 49, 2064 (1927). The resulting solution was acidified and extracted with other. The other solution was evapcrated; the residue distilled to isolate the e-mercaptobutyric acid. .

Subsequent treatment was the same as in Pramples XIV, part B, to yield dibutyltin S,S' bis(trimethylhexyl alpha mercaptobutyrate).

EXAMPLE XVII

The process of Example XIV was repeated using one mole of alpha bromovaleric acid in place of alpha chloropropionic acid. Dibutyltin S.S' bis(trimethylheryl alpha mercaptovalerate) resulted.

Alpha bromocaproic acid was prepared by the Hell-Vollhardt-Zelinsky bromination of 2 moles of caproic

One mole of the e-bromocaproic acid was used in the process of Example XVI to yield ultimately dibutyltin S.S' bis (trimethylhexyl e-mercaptocaproate).

acid.

EXAMPLE XIX

Two moles of caprylic acid were brominated by the Hell-Vollhardt-Zelimky method to yield a-bromocaptylic

One mole of the resulting e-bromocaprylic acid was used in the process of Example XVI, with the use of butyl alcohol in place of the 3,5,5 trimethylheranol, to yield dibutyltin S,S' bis(butyl e-mercaptocaprylate).

EXAMPLE XX

Ethyl e-mercapto-e-ethylcaproats was prepared by treating one mole of the ethyl e-bromo-e-ethylcaproste with two moles of 70% sodium sulfhydrate in 200 ml. of buryl alcohol saturated with hydrogen sulfide for 4-8 hours at reflux. The resulting ethyl s-mercapito-s-ethyl-caproate was treated with its equivalent of dibutylin oxide in refluxing toluene to yield, after removal of water and solvent, dibutyltin S,S' bis(ethyl e-mercapto-eethylcaproate).

EXAMPLE XXI

Capric acid (one mole) was treated by refluxing for one hour in thionyl chloride (200 ml.). To the resulting solution was then added 1.05 moles of bromine over a period of 2-3 hours. The mixture was allowed to stand overnight, the excess bromine and thionyl chloride removed by vacuum stripping. The residue was added to water to effect hydrolysis of the e-bromocapryl chloride. This is a modification of the process of Schwenk and Papa, J. Am. Chem. Soc. 70, 3627 (1948). The alpha- 40 bromo-capric acid was recovered and treated by the process of Example XIV to yield dibutyltin S.S. bis(butyl «-mercaptocaproate).

EXAMPLE XXII

One mole of pelargonic acid was treated by the process of Schwenk and Papa (J. Am. Chem. Soc. 70, 3627 (1948)), to yield ethyl a-bromopelargonate.

Using the process of Example XVI the ethyl e-bromopelargonate was converted ultimately to dibutyltin S,S' 50 bis(butyl e-mercaptopelargonate).

EXAMPLE XXIII

Lauric acid was treated by the process of Example XXI to yield dibutyltin S,S' bis(butyl e-mercaptolaurate).

EXAMPLE XXIV

One mole of palmitic acid was brominated by the standard Hell-Vollhardt-Zelinsky method. The e-bromopalmitic acid was treated by the method of Example XVI to yield dibutyltin S,S' bis(butyl a-mercaptopalmi-

EXAMPLE XXV

Two moles of stearic acid was treated by the method of Example XXII to yield dibutyitin S.S' bis(butyl smercaptostearate).

EXAMPLE XXVI

Two moles of phenylacetic acid were treated by the method of Example XXII to yield ethyl a-bromophenylacetate. The ethyl s-bromophenylacetate was treated with 2-mole equivalents of sodium sulfhydrate in 600 ml. of turn was treated with dibutyltin oxide in boiling toluene to

yield dibutyltin S,S' bis(ethyl o-mercaptophenylacerate). **EXAMPLE XXVII**

One male of diphenylacetic acid was treated by the process of Example XXVI. The intermediate ethyl ebromodiphenylacetate was a bluish green liquid which slowly became yellowish in color after standing. Dibutyltin S,S' bis(ethyl e-mercapto diphenylacetate) was the product obtained.

EXAMPLE XXVIII

Diethylacetic acid (2 moles) was treated by the process of Example XXII to yield dibutylin S.S' bis(trimethylhexyl e-mercaptodiethylacetate).

EXAMPLE XXIX

Water (1000 grams), 1,3 dibromopropane (2.50 moles, 502 grams) and phenol (2 moles, 188 grams) were stirred and heated to refinx and a solution of sodium hydroxide (1.9 moles, 76 grams) in 250 ml. of water was added slowly over a period of two hours. The mixture was then refluxed for 8 hours. The lower layer was separated, washed with a solution of sodium hydroxide, then with water, and dried over anhydrous sodium sulfate. It was then vacuum distilled to yield 3 phenoxypropyl bromide. This is a modification of a process given in J. Am. Chem. Soc. 71, 3163 (1949).

The 3-phenoxypropyl bromide was then added to an equivalent quantity of sodiomalonic ester in reflexing absolute alcohol sufficient to keep the sodiomalonic ester in solution. The sodiomalonic enter was prepared by dissolving an equivalent quantity of sodium in absolute ethanol (dried by the use of magnesium methoxide) and adding ethyl malonate to the sodium ethoxide solution. Both these reactions were carried out with suitable stirring. After the phenoxypropyl bromide had been added, refluxing and stirring were continued for two to three The solution was filtered, and the filtrate was distilled to remove ethanol and finally vacuum distilled to recover the diethyl 3-phenoxypropylmalonate.

The diethyl 3-phenoxypropylmalonate was refluxed with hydrobromic acid (200 mL of 48% HBr per mole of ester) in such a way as to remove the ethyl alcohol and ethyl bromide fromed. More hydrobromic acid was added as necessary. An exit tube from the top of the reflux condenser dipped under the surface of water in a small container served to indicate the rate at which carbon dioxide was evolved from decarboxylation of the substitoted malonic acid.

The 5-bromovaleric acid formed in this reaction was separated by dissolving in sodium bicarbonate solution, extracting with ether to remove phenol, acidifying the solution and finally extracting with ether. The ether solution was distilled to remove ether and the residue vacuum distilled to recover 5-bromovaleric acid.

Thiourea (2 mole equivalents) and the 5-bromovaleric acid were refluxed with sufficient water to dissolve the thiourea (about 100 ml./2 moles of thiourea) for 2 to 3 hours. Sodium hydroxide (5 moles/mole of 5-bromovaleric acid in 300 ml. of water) was then added and the solution refluxed for 20 hours. Acidification, ether extraction, and distillation of the ether extract yielded 5-mercaptovaleric acid.

Repeating the process in part B of Example XIV using: the 5-mercaptovaleric acid yielded dibutyitin S,S' bis-(trimethylhexyl 5-mercaptovalerate).

EXAMPLE XXX

The process of Example XXIX repeated where the 1,3 dibromopropane was replaced by 1.4 dibromobutane yieldwater to yield ethyl e-mercaptophenylacetate. This in 75 ed dibutyltin S.S' bis(trimethylhexyl 6-mercaptocaprosta).

EXAMPLE XXXI

Pentamethylene dibromide used in the process of Example XXIX yielded dibutyltin S,S' bis(trimethylhexyl 7-mercaptohexanoate).

EXAMPLE XXXII

Hexamethylene dibromide used in the process of Example XXIX (with the substitution of butyl alcohol for 3,5,5 trimethylhexanol) yielded dibutyltin S,S' bis(butyl 10 8-mercaptocaprylate).

EXAMPLE XXXIII

Decamethylene dibromide used in the process of Example XXIX, with the further substitution of butanol for the trimethylhexanol yielded dibutyltin S,S' bis(butyl 12-mercaptolaurate).

EXAMPLE XXXIV

Triethyl tin chloride (0.1 mole), decyl mercaptoacetate (0.1 mole) and toluene (100 ml.) were admixed. Sodium carbonate (0.05 mole) powder, anhydrous, was added gradually to the stirred solution. Carbon dioxide was evolved. Water of reaction was removed by refluxing with a moisture trap. The solution was filtered to remove sodium chloride. The toluene was stripped in vacuo. The pot residue, a liquid, was triethyl tin S(decyl mercaptoacetate). The material analyzed 25,43% Sn and 6.32% S.

EXAMPLE XXXV

Tributyl tin hydroxide (0.1 mole), mercaptoacetic acid (0.1 mole) and toluene (100 ml.) were admixed. Water of reaction was removed by refluxing with a moisture 85 trap. The toluene was stripped in vacuo. The resulting material, a viscous liquid, was tributyl tin S(mercaptoacetic acid). It analyzed 31.36% Sn and 9.35% S.

EXAMPLE XXXVI

Triethyl tin chloride (0.1 mole), mercaptoacetic acid (0.1 mole) and 100 ml. of toluene were admixed. Sodium carbonate (0.05 mole), anhydrous, was added slowly to the stirred solution. Carbon dioxide was evolved. Water of reaction was removed by means of a water trap. The sodium chloride was removed by filtration. The toluene was stripped in vacuo. The resulting viscous liquid was triethyl tin S(mercaptoacetic acid). It analyzed 38.21% Sn and 12.12% S.

EXAMPLE XXXVII

Butyl tin trichloride (0.1 mole), mercaptoacetic acid (0.3 mole) and toluene (100 ml.) were admixed. Sodium carbonate, anhydrous powder (0.15 mole), was added gradually to the stirred solution. Carbon dioxide was

evolved. Water of reaction was removed by refluxing with a moisture trap. Sodium chloride was removed by filtration. The toluene was removed in vacuo. The resulting solid was butyl tin S.S.S" tris(mercaptoacetic acid). It analyzed 20.90 Sn and 17.29 S.

It has also been discovered and is a feature of this invention that the novel organo-tin derivatives of mercapto acids and mercapto acid esters will function as excellent stabilizers for halogen containing resin compositions, particularly vinyl halide resin compositions containing plasticizers, and when intimately dispersed therein, will provide plastic compositions of improved resistance to heat deterioration. Excellent films are obtained from the aforesaid plastic compositions which exhibit a high degree of stability. The optimum concentration of the novel organo-tin derivatives, useful as a stabilizer is between 0.5-10% based on the weight of the vinyl resin. The resin compositions containing this concentration of stabilizer produce a stable and colorless plastic film. which remains clear at elevated temperatures as evidenced by the results of the heat tests in Table I.

Polyvinyl chloride resins and copolymer resins were employed in these tests. The stabilizer was incorporated into a mixture of 100 parts by weight of resin and 35-50 parts by weight of a plasticizer, or a mixture of plasticizers. The weight of the stabilizer was determined by its analysis. Thus enough stabilizer was used to provide the same weight of combined tin as provided by two parts of dibutyl-tin S,S bis(nonyl mercaptoacetate) (0.374 part of tin by weight). This was done in order to have a means of determining the relative merits of the various compounds.

The mixture was then milled for 5 minutes on a two-roll differential speed mill heated to 320 to 325° F., and removed as a sheet. Portions of the sheet were then placed in a single cavity mold (6" by 6" by 40 mils), and preheated to 275° F. The mold was placed on a Preco press and raised to 320° F. under 10,000 pounds total pressure. When the mold reached 320° F. the pressure was increased to 40,000 pounds and held until the temperature reached 330° F. This procedure required five to five and one-half minutes. The mold and press platens were then cooled and the pressed sheet removed.

The pressed sheet was cut into one inch by six-inch strips and these placed in clips on a tray so that the strips would hang vertically. The tray was then placed in a circulating air oven held at 320° F. Samples were recovered after one hour, two hours, three hours, and four hours of heat aging. Samples were rated visually, the degree of stability of the vinyl resin film being represented by the depth of the colorations, wherein C is colorless, VSY is very slight yellowing, SY is slight yellowing, Y is yellowed, OY is orange yellow, rbr is reddish brown, br is brown, and bl is black.

TARLE I

| Composition . | | | Appearance after heat aging | | | |
|---------------|--|---|-----------------------------|---------------|----------|--------------------------------|
| Stabilizar | 100 parts rests. | Plasticier | 1 hr. | 2 hr. | 8 hr. | 4 hr. |
| None | Geon 101 2 parts/100 parts Geon 101 restn. | 50 parts Fiscal DOP 50 parts Fiscal DOP | ₩ | pt | u | . • |
| From Br. L | Geon 101 EP (from B. P. Goodrich Co.; polyvinyi chloride resin, homo- polymer). | 50 parts Florel DOP (di- cotyl phthalate). | 0 | C | 0 | VBY. |
| From Er. 2 | Same Same Geon 101—polyvinyi chlo- rida, resin, homopoly- mar; particle star; 100% | Same | C | Stable (grey) | C | VSY. VSY. Stable (grey). |
| | through 63 mesh acrem, specific gravity L61, average sortime extract 13%, average specific viscosity 0.45 (at 20° O. of 0.4% solution in ni- tros benume). | · •. | <i>:</i> . | | : | |

TABLE I-Outland

| Company of the control of the contro | | | | | | |
|--|--|--|-----------------------------|-------------|---------------|---------------|
| Stabilizar | Competition | | Approximen other hant aging | | | |
| 11432 | 200 parte sente | Platities . | 1 le. | S he. | 8 kg. | 4 hr. |
| Press Ex. L | Gen 201 EP | | . 0 | 0 | | |
| Do | Oess 165 E7 | - 8000 | | (100) older | C. C. (C.A.) | Stable (C-G). |
| From Ex. C | Geim 181 | - | (FT) | C. (5-47) | C | J'VDI. |
| From No. 7 | Occa 101 EP | | el Vermaner | O | (1973) attack | Stable (com). |
| From Ex A | Ultren 200 Odersta | | - g | Q | 10 | VBY. |
| | Chambles On restrict | | . C | C | C | VEZ. |
| From Ex. 10. | views state the restal. | Berne | 1_ | | · | • |
| From Ex. 11. 2 parts dibuty! tts. 8, 9 | . Same | The same of the sa | - <u>S</u> | C. | Ç | YEX. |
| pe (non) member | Ultren 200 | . 40 perts Flexal DOP, 16 | 8 | O | Tex | PDF. |
| acatata). | 1 | 60 perts Floral DOP, 16 perts 8-160 (gray) di- phenyi phesphate- Momental. | 1 | - | 0 | VEX. |
| Da | - Same | Moneante). | l | | | • |
| Da | | - Wind Dur, Street | C | 0 | VEY | 8T. |
| ~ | | D parts DOP, 30 parts \$- | 0 | 0 | VOY | |
| Da | Same | 300 | | | ABX | SY. |
| Do | | 10 parts DOP, 40 parts 8- | C | 0 | 6T | T. |
| /// | Geon ittl EP (pokyvinyl chloride ruin, kome | 40 parts DOP, 10 parts Lindol (tricrenyl phos- phate, Columns Corp. | 0 | 0 | C | |
| | DOLLARS, TAIL, PARTY | Linda (tricrity) pho- | | | V | C. |
| De | | of Art.) | ! · | | ı | |
| | | of Ass.). 20 parts DOP, 30 parts. | 0 | a | 0 | |
| De | Barna | Lindel Don | | | 0 | VET. |
| Da | Same | 20 parts DOP, 20 parts | 0 | 0 | VBY | VEY. |
| | | 10 parts DOP, 40 parts | 0 | 0 | V87 | VEY. |
| Da | Seme | I GO THEFTO TO COME IN MARKET CO | | | | VOI. |
| | | 141 (an ary) alky) phos- phase-Manageria). 20 parts DOP, 20 parts 8- | C | C | O) | C. |
| De | Same | phets Monanto). | | i | 1 | |
| Do | | | O | C | C | VEX. |
| | E | 20 parts DOP, 20 parts 8- | 0 | 0 | اـــــا | |
| Do | Same | I 1601. I | | | | VEY. |
| 2.5 pairts (Dipty) ttn. 6.8' | | 16 parts DOP, 40 parts 8- | O | C | VSY | 5 7. |
| pre (DOM) EDECATOR | CASE AND CONTACT OF | 40 parts DOP | 0 | 0 | a | a |
| acetata). | Chicago colpolymer re- | | | | - | u. |
| Da | Occa 322 (polyviny) chlo- ride - polyviny) idease chloride expolymer sa- dia. B. F. Occarin Col. V2-10 (Namparett Cham. Col.). A (polyviny) chlo- ride saim, Goedyner Thr & Bubbar Col. | | - 1 | i | 1 | |
| | VA-W Cressponds Chan. | 85 perts DOP | 0 | o | al | a . |
| Do | Priorie & (petrylant dila. | S parts DOP | VEX | | | |
| | Title rects, Goodyness | | T= 4 | VEY | 5 T | T. · · |
| Da | U-SOL | Street DOD 15 | _ 1 | _ 1 | · i | <i>:</i> |
| Da | | 25 parts DOP, 15 parts \$- | 0 | C(| O | VEY. |
| Do | Georg 1/8 | # parts DOP | Light gray | Light grey | 4 | |
| Do | QYNA (polyviny) ablo- | Seme | C | U | 5 | Held grey. |
| l | ride-polyvinyl sertate | | · | 76Y | 17 | r. |
| 2 perts dibuty! the di- | QYNA (polyviny) chlo- ride-polyviny) certain captlymer Bekelten). Geom itt | STORES TOOL OF THE PARTY OF THE | _ 1 | | · 1 | |
| lattrate. | 4- | XO | Y(| DYYO | | |
| | | | Y | DT | - | |
| 3 parts dispty! tin malests. | | Sema : | | 1 | | • . |
| 3 parts dibuty) the dilem- | | 20 parts DOP, 20 parts Lindel | OT | 757 | br | · . |
| | Geon 101 | | 1 | | | • |
| 1.35 parts. Ex. 34 | Geon M. Vitron 800 | D parts DOP | Ç <u></u> <u>1</u> | (BY | Y | T. |
| 0.46 perts. Ex. St. 1.6 perts, Ex. ST. | do | | Y | 1 | <u> </u> | E |
| | | | L 100 | | | . |
| This table clearly | discloses the | | | | | =• |

This table clearly discloses the unusual stabilizing activity of the novel mercapto-acid ester derivatives of organo tin compounds in comparison to a resin without a stabilizer and with known stabilizers. A colorless film evidences the highest degree of stability. This superior stabilizing effect is particularly evident in polyvinyl chloride rusin compositions containing phosphate-type plasticizers, the use of which has heretofore been limited due to their instability. The novel organo-tin compounds of the present investion impart sufficient stabilization to such compositions to permit making stable fireproof plastic articles therefrom.

Is order to test the effectiveness of the compounds of this invention as light stabilizers, films of 5 mil thickness were prepared by the same methods described above. In each case dibutythin S.S' bis(nony) mercaptoacetate) was used as a stabilizer. These films were exposed in an Atlas Electric. Devices Company Fadoometer FDA-2 and examined daily until failure occurred. The compositions of the films, the time at which failure was first noted and the type of failure are given in Table II. Type of failure is indicated by a code in which A represents sweating, B specking or spotting, S stiffening and E embrittlement. It should be noted that up to 1000 hours no stiffening or embrittlement had occurred.

TABLE II

| • | | | | | | | | |
|----------------|--|---|--|---------------------|--|--|--|--|
| : | 70: | | | | | | | |
| 55 | Restn | Plasticier | Fading hours | Sheet, | | | | |
| 60 65 70 | 100 parts polyvinyl chloride ratin (Geos III EP). Do. Do. 100 parts polyvinyl chloride ratin (U-200). Do. Do. Do. Do. Do. Do. Do. D | 60 parts of disctyl phrhalate (DOP), 10 parts of an aryl altyl phrephate (8-141). 30 parts DOP, 30 parts 8-141. 30 parts DOP, 40 parts 8-141. 10 parts DOP, 40 parts 8-141. 15 parts DOP, 15 parts 8-141, 1 part dibutylin 8.8" 141, 1 part dibutylin 8.8" | 1 00 1 00 1 00 1 00 1 00 1 00 1 00 1 0 | 4 43434343 43 43 43 | | | | |
| | | - tata. | | • | | | | |

Other uses of the organo-tin-mercaptoscid and mercapto acid ester derivatives are as stabilizers for other 13

chlorinated materials, as rubber accelerators, rubber antioxidants, lube oil additives and polymerization accelerators.

While the invention has been described with reference to various examples and embodiments, it will be apparent to those skilled in the art that various modifications may be made, and equivalents substituted therefor, without departing from the principles and true nature of the present invention.

Having described the invention, what is desired to be se- 10 cured by Letters Patent is:

1. A compound having the structural formula

R_sSnSR'COOR"

wherein R is selected from the group consisting of alkyl, 15 aryl and aralkyl radicals, R" is from the group consisting of hydrogen and of an alkyl, aryl, alkaryl and aralkyl radicals, and R' is selected from the group consisting of alkylene and aralkylene radicals.

2. As a composition of matter, a trihydrocarbon tin 20 derivative of an aliphatic mercapto monocarboxylic acid ester containing the group

3. A method of preparing compounds as defined in claim 2 which comprises reacting a trihydrocarbon substituted compound of the class consisting of oxides, hydroxides and halides of tin and stannoic acid with a

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compound having the formula HSR'COOR" in which R" is selected from the group consisting of hydrogen, alkyl, aryl and aralkyl radicals, and R' is selected from the group consisting of alkylene and aralkylene radicals.

- 4. Tributyltin S(decyl mercaptoacetate).
- 5. Triphenyitin S(decyl mercaptoacetate).
- 6. Triethyl tin S(decyl mercaptoacetate).
- 7. Tributyl tin S(mercaptoacetic acid).

fined in claim 1.

- 8. Triethyl tin S(mercaptoacetic acid).
 9. A stabilizer for halogen-containing resins and plas-
- ticizers of the character defined in claim 1.

 10. A composition comprising a halogen-containing resin stabilized with a compound of the character de-
- 11. A composition comprising a halogen-containing resin stabilized with about 1/2 to 10% by weight of a compound of the character defined in claim 1.
- 12. A halogen containing resin having intimately dispersed therein a trihydrocarbon tin derivative of an aliphatic mercapto monocarboxylic acid ester containing the group

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